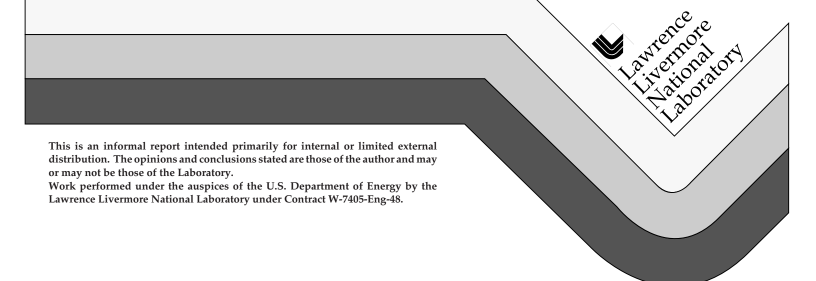
¹²⁹I Interlaboratory Comparison: Phase I and Phase II Results

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July 1997



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UNITED STATES MEMBER STATE SUPPORT PROGRAM TO IAEA SAFEGUARDS

DEPARTMENT OF ENERGY DEPARTMENT OF STATE ARMS CONTROL AND DISARMAMENT AGENCY NUCLEAR REGULATORY COMMISSION DEPARTMENT OF DEFENSE

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July, 1997

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¹²⁹I Interlaboratory Comparison: Phase I and Phase II Results

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Abstract

An interlaboratory comparison exercise for ¹²⁹I was organized and conducted. A total of nine laboratories participated in the exercise to either a full or limited extent. In Phase I of the comparison, a suite of 11 samples were measured. The suite of samples contained both synthetic 'standard type' materials (i.e., AgI) and environmental materials. The isotopic ¹²⁹I/¹²⁷I ratios of the samples varied from 10⁻⁸ to 10⁻¹⁴. In this phase, each laboratory was responsible for its own chemical preparation of the environmental samples. The ¹²⁹I AMS measurements obtained at different laboratories for prepared AgI were in good agreement. However, large discrepancies were seen in ¹²⁹I AMS measurements of environmental samples.

Because of the large discrepancies seen in the Phase I intercomparison, a subsequent study was conducted. In Phase II of the comparison, AgI was prepared from two environmental samples (IAEA 375 soil and maples leaves) by three separate laboratories. Each laboratory used its own chemical preparation method with each of the methods being distinctly different. The resulting six samples (two sets of three) were then re-distributed to the participating ¹²⁹I AMS facilities and ¹²⁹I/¹²⁷I ratios measured. Results and discussion of both the Phase I and Phase II interlaboratory comparison are presented.

This work was done as part of U.S. Support Program Task DOE.10 (Special Analysis Procedures for Detecting Undeclared Activities).

We wish to thank the U.S. Department of Energy, International Safeguards Division, NN-44 for their support of this project.

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Introduction

In April 1993, at an International Atomic Energy Agency (IAEA) consultant's meeting in Vienna, several technical issues relating to environmental monitoring, sampling, and analyses for the detection of undeclared nuclear activities were raised and discussed. In particular, Accelerator Mass Spectrometry (AMS) was identified as an important high sensitivity analysis technique for the detection of ¹⁴C and ¹²⁹I. Detection of ¹⁴C using AMS is a well established technique used in a wide range of scientific applications. About thirty laboratories worldwide have established a ¹⁴C AMS analysis capability. Through the efforts of the radiocarbon dating community, appropriate standards, blanks, and analysis protocols have been determined and numerous interlaboratory comparisons have been performed. Given established sample handling and preparation procedures, the precision and accuracy of a ¹⁴C AMS analysis is seldom in doubt. Unfortunately, this degree of reliability has not yet been achieved for ¹²⁹I AMS analysis. A very limited number of laboratories have established analysis techniques for ¹²⁹I AMS, and typically, each laboratory has its own independent standards and blanks by which abundance concentrations are determined.

Because of these unresolved questions about ¹²⁹I AMS measurements, Lawrence Livermore National Laboratory (LLNL) proposed and was funded by the U.S. Department of Energy to sponsor an ¹²⁹I intercomparison exercise on behalf of the IAEA. From the IAEA's point of view, the purpose of the ¹²⁹I intercomparison exercise was to assess the suitability and effectiveness of ¹²⁹I AMS measurements for possible safeguards use. Detection of ¹²⁹I is important because the IAEA has identified ¹²⁹I as a potential signature of reactor or reprocessing operations. For the ¹²⁹I intercomparison exercise to be of maximum usefulness to the IAEA, it was important that the exercise contain environmental material of the type that the IAEA would expect to acquire on a typical field trial or inspection. As described by the IAEA, types of environmental samples that might be acquired on a field trial include swipes, filters, soils, grasses, lichens or moss, deciduous leaves, tree bark, pine needles, sediments, water, and water biota (e.g., algae, mussels, plants).

Eleven laboratories were invited to take part in the exercise and, at the time of writing, a total of nine laboratories participated in the exercise to either a full or limited extent. Laboratories that participated in the exercise are listed in the acknowledgments.

Phase I Samples

In February of 1995, LLNL prepared and shipped a suite of 11 ¹²⁹I intercomparision samples. The suite of samples was developed from discussions with the IAEA and contained both synthetic 'standard type' materials (e.g., AgI) and environmental materials of the type that the IAEA would expect to acquire on a typical field trial or inspection. The specific samples were:

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Prepared AgI. ^{129}I/^{127}I ratio calculated to be 90308 x 10^{-15}.
Sample #1:
                      Prepared AgI. ^{129}I/^{127}I ratio calculated to be 45474 x 10^{-15}.
Sample #2:
                     Prepared AgI. ^{129}I/^{127}I ratio calculated to be 21729 x 10^{-15}.
Sample #3:
                      Prepared AgI. ^{129}I/^{127}I ratio calculated to be 4922 x 10^{-15}.
Sample #4:
                      Water sample. <sup>129</sup>I/<sup>127</sup>I ratio unknown but less than 10<sup>-10</sup>.
Sample #5:
                      Spiked swipe. A Whatman filter paper spiked with <sup>129</sup>I.
Sample #6:
                     Pine needles. <sup>129</sup>I/<sup>127</sup>I ratio unknown but less than 10<sup>-10</sup>.
Sample #7:
                     Maple leaves. <sup>129</sup>I/<sup>127</sup>I ratio unknown but less than 10<sup>-8</sup>.
Sample #8:
                      Dried sea weed. <sup>129</sup>I/<sup>127</sup>I ratio unknown but less than 10<sup>-8</sup>.
Sample #9:
                      Soil. <sup>129</sup>I/<sup>127</sup>I ratio unknown but less than 10<sup>-10</sup>.
Sample #10:
                     Woodward Iodine. ^{129}I/^{127}I ratio approximately 50 x 10^{-15}.
Sample #11:
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As can be seen, the isotopic $^{129}I/^{127}I$ ratio of the samples varied from 10^{-8} to 10^{-14} .

The first three AgI samples had ¹²⁹I/¹²⁷I ratios that are comfortably measured by the AMS technique. These three samples provided some statistically meaningful intercomparison of the AMS technique at the various participating laboratories. The fourth AgI sample had an ¹²⁹I/¹²⁷I ratio much lower than the first three AgI samples and provided a 'low-level' intercomparison sample. The Woodward iodine sample was included to help in the determination of backgrounds. All prepared AgI samples were precipitated from a bulk solution that was derived by successive dilutions of a National Institute of Standards and Technology (NIST) standard material. The ¹²⁹I/¹²⁷I ratio of the original NIST standard material was 0.4091.

The water sample was created using de-ionized purified water. Iodine was added to this water in the form of potassium iodide and, to prevent loss of iodide, both NaOH and sodium bisulfite were added. The iodine concentration for this sample was approximately 250 μg I /g of H_2O .

The remaining five samples on the list were included to represent environmental samples that the IAEA would typically collect. The swipe sample was a Whatman filter paper spiked with a known amount of ¹²⁹I. The pine needle and soil samples were collected in the vicinity of LLNL and were believed to have ¹²⁹I concentrations such that the high sensitivity of the AMS technique is truly required to obtain isotopic abundances. The seaweed and maple leaf samples were not collected locally to the LLNL area and possessed sufficiently high ¹²⁹I concentrations that they could be measured by both thermal emission mass spectrometry and AMS. The seaweed, maple leaf, pine needle, and soil samples provided to the participating laboratories were representative aliquots from a larger supply of sample material. To ensure homogeneity of the samples, each sample was extensively ground and mixed. All samples were prepared identically, at the same time, and under the same conditions.

It is also important to note that the environmental samples included in the ¹²⁹I intercomparison exercise were never intended to become 'NIST type' environmental standards. Our main intent with the chosen set of environmental samples was to help the IAEA determine to what level one can expect agreement between results obtained from different ¹²⁹I AMS laboratories.

Phase I Intercomparison Procedure

The suite of samples were distributed to all laboratories expressing an interest in taking part in the ¹²⁹I AMS intercomparison. The only information given to the participating laboratories regarding the ¹²⁹I/¹²⁷I ratios of the individual samples was an approximate guide to the upper limit of the expected ¹²⁹I/¹²⁷I ratio. Sufficient sample material was given to each laboratory to allow several repeat measurements. Laboratories were asked to report the results for the AgI samples (samples 1, 2, 3, and 4) and the Woodward Iodine sample (sample 11) as ratios (i.e., number of ¹²⁹I atoms per number of ¹²⁷I atoms). For the water, swipe, and other environmental samples (samples 5, 6, 7, 8, 9, and 10), laboratories were asked to report results as concentrations (i.e., the number of ¹²⁹I atoms per gram of sample).

Phase I Results

The results of the Phase I 129 I intercomparison are shown in Table 1 and in Figures 1 and 2. To preserve the anonymity promised to the participating

laboratories, individual laboratories are identified only by code. Because some of the participants in the exercise did not have the requisite chemical preparation lines and procedures necessary to prepare and measure the environmental samples, some of the laboratories were only able to report results for the AgI samples.

As can be seen from Figure 1 and Table 1, the agreement between ¹²⁹I/¹²⁷I ratios from the different laboratories is excellent for the AgI samples (samples 1, 2, 3, and 4). Table 1 also lists the un-weighted means and standard deviations of the results from the AgI samples. For the AgI samples, the differences of the un-weighted means from the expected ¹²⁹I/¹²⁷I ratios were always less than 5%, while the standard deviations of the means were generally around 5%. This agreement of ¹²⁹I/¹²⁷I ratios for AgI samples between different laboratories is remarkable considering the difficulty of the analytical technique as well as the fact that different laboratories tend to use their own independent standards and blanks.

Looking at Figure 2 and Table 1, however, one can see that, except for sample 5, there is large disagreement between measured ¹²⁹I concentrations from the various environmental samples. For example, differences in ¹²⁹I concentrations of two to three orders of magnitude were reported for samples 8 and 9. These large differences between reported ¹²⁹I concentrations are particularly disappointing considering the excellent agreement obtained between the AgI samples. One possible cause of such disagreement was thought to be the differing chemical preparation methods used by each laboratory. These differing chemistries could contribute to differences in extraction of ¹²⁹I and may be an explanation for the widely differing results seen from environmental samples.

Another possible explanation for the widely differing results from environmental samples is ¹²⁹I contamination of the chemical preparation laboratories. While contamination was not believed to be a problem at all the participating laboratories, one of the laboratories suffered severe cross contamination problems from previous samples. This particular laboratory eventually solved the contamination problem by building a new chemical extraction line in a different building..

¹²⁹I Workshop

On May 16, 1996, in conjunction with the 7th International Conference on Accelerator Mass Spectrometry, LLNL hosted a one day pre-conference workshop that concentrated on the application of the ¹²⁹I AMS technique. Most of the participants in the ¹²⁹I intercomparison exercise were present at the workshop. A major part of this workshop was the discussion of results from the present ¹²⁹I intercomparison exercise, ¹²⁹I sample preparation methods, and possible tests needed to understand the large discrepancies in ¹²9I results obtained for the environmental samples. As was expressed at both formal and informal discussions, widely differing ¹²⁹I environmental results could possibly be explained by differences in ¹²⁹I extraction chemistry. To better characterize this extraction chemistry theory, it was agreed to conduct a Phase II of the ¹²⁹I intercomparison exercise.

In Phase II of the ¹²⁹I intercomparison exercise, it was agreed that AgI was to be prepared from two different environmental samples by three separate laboratories. The resulting six samples (two sets of three) were then to be re-distributed to all the ¹²⁹I AMS facilities and ¹²⁹I/¹²⁷I ratios measured. It was believed that results from this Phase II of the exercise would help determine whether the large discrepancies seen in results obtained for the Phase I environmental samples was due to chemical extraction procedures or due to some problem with the analytical method.

Phase II Samples

Three laboratories volunteered to chemically prepare samples for Phase II of the ¹²⁹I intercomparision. The three laboratories were 1) Texas A&M University, College Station, Texas, 2) PSI/ETH, Zürich, Switzerland, and 3) LLNL, Livermore, California. Each laboratory uses its own ¹²⁹I chemical preparation method. Texas A&M prepares ¹²⁹I samples using an alkali leach and fusion method^[1,2]. PSI/ETH works in conjunction with ZSR (Center of Radiation Protection and Radioecology) at the University of Hannover, Germany and uses a dry-ash combustion procedure to prepare environmental ¹²⁹I samples^[3]. LLNL uses an wet-ash distillation method to prepare environmental ¹²⁹I samples.

In June of 1996, LLNL shipped two environmental samples and potassium iodide carrier solution to three laboratories listed above. The two environmental samples were soil and maple leaves. The soil used in Phase II of the ¹²⁹I intercomparision was IAEA #375 reference material (IAEA reference sheet, reference material IAEA 375-soil, reference number G4.12). The soil is reported to

have an ^{129}I concentration of 1.7(±0.4) x 10^{-3} Bq/kg or 1.2(±0.3) x 10^{9} ^{129}I atoms/gram material. Approximately 10 grams of soil was provided to each of the three chemical preparation laboratories.

The maple leaves used in Phase II of the 129 I intercomparision were collected near the town of Sequim, Olympic Peninsula, Washington State, USA. These leaves were also used as sample 8 in Phase I of the intercomparison. The precise 129 I concentration of the maple leaves is unknown but was tentatively measured in Phase I of the intercomparison exercise to be between 3.1×10^7 and 5.2×10^9 129 I atoms/gram material. Approximately 9 grams of leaves were provided to each chemical preparation laboratory.

In conjunction with the soil and maple leaves, LLNL also provided KI carrier solution to each of the chemical preparation laboratories. The KI Carrier Solution was a 1 Normal or 12.3 mg I/gram solution. The 129 I/ 127 I ratio of KI precipitated from the carrier solution has been measured to be less than 2 x 10^{-14} .

Each chemical preparation laboratory was asked to make approximately 50 to 60 mg of AgI from both the soil and leaf samples. This amount of AgI was enough to provide each of the ¹²⁹I AMS measurement laboratories with approximately 5 to 6 mg of AgI. Each laboratory was asked to add enough carrier to each sample so that one could make the assumption that all the ¹²⁷I comes from the carrier and all the ¹²⁹I comes from the sample.

In addition to the six environmental samples, LLNL prepared an AgI sample that was precipitated from a bulk solution that had been derived by successive dilution's of a NIST standard material. This sample was identical to sample 2 used in Phase I of the intercomparison. This sample is useful to check for normalization errors and to provide a statistically meaningful comparison of the AMS technique at the various participating laboratories using a well constrained sample.

Details of the seven individual samples are as follows:

Sample 31. AgI prepared from NIST standard material Same as sample 2 used in Phase I of the intercomparison $^{129}\text{I}/^{127}\text{I}$ ratio calculated to be 45474 x 10^{-15}

Sample 32. AgI prepared from IAEA #375 reference material soil

Amount of sample used: 2.84 g

Amount of KI carrier added: 69.7 mg

Sample 33. AgI prepared from maple leaves

Amount of sample used: 5.30 g

Amount of KI carrier added: 100 mg

Sample 34. AgI prepared from IAEA #375 reference material soil

Amount of sample used: 9.98 g

Amount of KI carrier added: 29.89 mg

Sample 35. AgI prepared from maple leaves

Amount of sample used: 5.87 g

Amount of KI carrier added: 22.76 mg

Sample 36. AgI prepared from IAEA #375 reference material soil

Amount of sample used: 1.20 g

Amount of KI carrier added: 61.5 mg

Sample 37. AgI prepared from maple leaves

Amount of sample used: 3.60 g

Amount of KI carrier added: 61.5 mg

Phase II Intercomparison Procedure

By September of 1996, LLNL had received bulk aliquots of AgI from the three chemical preparation laboratories. The bulk AgI was subdivided at LLNL and distributed to all laboratories that expressed a willingness to take part in Phase II of the ¹²⁹I AMS intercomparison exercise. The only information given to the participating laboratories regarding ¹²⁹I/¹²⁷I ratios of the individual samples was an approximate guide to the upper limit of the expected ¹²⁹I/¹²⁷I ratio. All participating laboratories were asked to report results as ratios (i.e., number of ¹²⁹I atoms per number of ¹²⁷I atoms).

Phase II Results

Results of the Phase II ¹²⁹I intercomparison are shown in Figures 3, 4, 5, 6, and 7 and in Tables 2 and 3. In order to preserve the anonymity promised to the participating laboratories, individual laboratories are identified only by code.

With the exception of one measurement, the agreement between ¹²⁹I/¹²⁷I ratios from different laboratories is good for sample 31 (Figure 3 and Table 2).

Including laboratory B, the difference of the un-weighted mean from the expected ¹²⁹I/¹²⁷I ratio is about 10% with a standard deviation of the means of about 15%. Excluding laboratory B, the difference of the un-weighted mean from the expected ¹²⁹I/¹²⁷I ratio is approximately 5% with a standard deviation of the means of about 4%. As was the case in Phase I of the ¹²⁹I intercomparision, the agreement of ¹²⁹I/¹²⁷I ratios for the AgI 'standard type' material is excellent.

Results from the IAEA #375 reference material soil are shown in Figures 4 and 5 and in Tables 2 and 3. Measured ¹²⁹I/¹²⁷I ratios for samples 32, 34, and 36 are shown in Figure 4 and in Table 2 while ¹²⁹I concentrations calculated from the measured ¹²⁹I/¹²⁷I ratios are shown in Figure 5 and in Table 3. The agreement of ¹²⁹I/¹²⁷I ratios amongst the various AgI aliquots is good (Figure 4). After ¹²⁹I/¹²⁷I ratios have been converted to ¹²⁹I concentrations, as is shown in Figure 5 and Table 3, the three differing chemical preparation methods give consistent results for the ¹²⁹I concentration of the IAEA #375 soil. Examining the un-weighted means or 'average' ¹²⁹I concentration, however, it appears that all three differing chemical preparation methods yield an average ¹²⁹I concentration that is slightly lower than the reported IAEA ¹²⁹I concentration value of 1.7(±0.4) x 10⁻³ Bq/kg or 1.2(±0.3) x 10⁹ ¹²⁹I atoms/gram material. It may be premature, however, to draw major conclusions from this slight systematic shift considering the 25% error in the reported IAEA concentration and the 10-20% standard deviations associated with the AMS results.

Results from the maple leaf sample are shown in Figures 6 and 7 and in Tables 2 and 3. Measured ¹²⁹I/¹²⁷I ratios for samples 33, 35, and 37 are shown in Figure 6 and in Table 2 while ¹²⁹I concentrations calculated from the measured ¹²⁹I/¹²⁷I ratios are shown in Figure 7 and in Table 3. As one can see from Figure 6 the agreement of ¹²⁹I/¹²⁷I ratios amongst the various AgI aliquots is poor relative to the agreement obtained for the soil samples. Because of the poor agreement of ¹²⁹I/¹²⁷I ratios, ¹²⁹I concentrations are also in poor agreement as is shown in Figure 7 and Table 3. Although there is a factor of four difference between the un-weighted mean ¹²⁹I concentration of samples 33 and sample 35 (with sample 37 lying somewhere in between), it is hard to draw any conclusions about the three differing chemical preparations due to the large standard deviations. What is clear, however, is that the different AMS laboratories cannot obtain the same ratio for AgI independent of the chemical preparation method.

Discussion of Phase II Results

Results of Phase II of the ¹²⁹I AMS intercomparison show good agreement of ¹²⁹I/¹²⁷I ratios for AgI 'standard type' material. This good agreement was also seen in Phase I of the intercomparison.

The phase II ¹²⁹I AMS intercomparison results show agreement with ¹²⁹I concentrations in IAEA #375 soil measured by other techniques. This agreement is independent of the chemical preparation method.

The phase II ¹²⁹I AMS intercomparison results show relatively poor agreement of ¹²⁹I concentrations in the maple leaves. This relatively poor agreement of ¹²⁹I concentrations appears largely to be due to the fact that different AMS laboratories do not obtain the same ratio from aliquots of the same AgI. We are unable at this time to explain the cause of these disagreements. We only offer the following comments:

1) Is there something in the maple leaves which results in an 'effective' contamination of the AgI, and which the various ¹²⁹I AMS measurement laboratories have trouble discriminating against? This contaminant must be common to all three chemical preparation methods. To test this hypothesis, we plan on performing Proton Induced X-Ray Emission (PIXE) measurements of the various samples to see if gross elemental differences between AgI made from the soil and AgI made from maple leaves can be detected.

Is there a problem with the ¹²⁹I AMS analytical technique? We note that, in general, laboratories N, V, and M measured a 'high' ¹²⁹I/¹²⁷I ratio for samples 33, 35, and 37 while Laboratories W, K, and Q measured a 'low' ¹²⁹I/¹²⁷I ratio for samples 33, 35, and 37. If we make the assumption that the individually distributed samples each have a uniform ¹²⁹I/¹²⁷I ratio, does this mean that laboratories N, V, and M have a problem with discriminating against some unknown and interfering nuclide? If there is a problem with background or an unknown nuclide, one might make the assumption that those laboratories with the highest terminal potentials and most sophisticated spectrometers would the best at discriminating against this supposed contaminant. We note, however, that some of the 'high' ¹²⁹I/¹²⁷I ratios come from laboratories with

the 'sophisticated' spectrometers while some of the 'low' ¹²⁹I/¹²⁷I ratios come from laboratories with the relatively 'simple' spectrometers.

- 2) Is there a problem with ¹²⁹I contamination at some point in the process? It has been suggested that, since the isotopic ratios of the soil samples are somewhat higher than the maple leave samples, a possible background correction is more severe for the maple leaves. Contamination is certainly a possible explanation for the disagreement seen. We note, however, that maple leaf samples 33 and 35 had un-weighted mean ratios that were not that much different than soil sample 36 and that sample 36 did not have the large variances in reported 129 I/127 I ratios that would be indicative of a contamination problem in the analytical method. If there is a contamination problem, we feel it has to be somewhat unique to the maple leave samples. We also feel that any potential contamination would have had to occur after chemical preparation of samples because there is not a statistically significant difference in the un-weighted mean 129 I concentrations of the maple leaf samples. If contamination of the maple leaf samples had occurred during chemical preparation we should have seen large differences in the un-weighted mean ¹²⁹I concentrations. Most likely, any contamination of the maple leaf samples would have had to occur at LLNL during sample redistribution or at the various ²⁹I AMS analytical laboratories.
- 3) When calculating the ¹²⁹I concentrations for samples 32 through 37, no adjustments were made for less than 100% iodine recovery. It was assumed that the carrier iodine and iodine from the sample were recovered in the same proportion. Iodine recoveries could well differ for the three different chemical preparation methods. While recovery could have an effect when comparing the average ¹²⁹I concentrations, iodine recovery differences do not explain the large standard deviations obtained from samples 33, 35 and 37.

Conclusions

Because of unresolved questions about ¹²⁹I AMS measurements, an interlaboratory comparison exercise for ¹²⁹I has been organized and conducted. The primary purpose of this ¹²⁹I intercomparison exercise was to assess the suitability and effectiveness of ¹²⁹I AMS measurements for possible IAEA safeguards use. From the data collected so for, it appears that:

- 1) Good agreement of ¹²⁹I/¹²⁷I ratios can be obtained for AgI 'standard type' material using AMS.
- 2) Good agreement of ¹²⁹I concentrations in soils (in this case IAEA #375 soil) can be obtained using AMS. This agreement can be obtained largely independent of the chemical preparation method.
- 3) Relatively poor agreement of ¹²⁹I concentrations in low activity organic material (in this case maple leaves) is obtained using AMS. The cause of this poor agreement is unknown. For this class of samples, more effort is needed to understand the cause of this large discrepancy.

Acknowledgments

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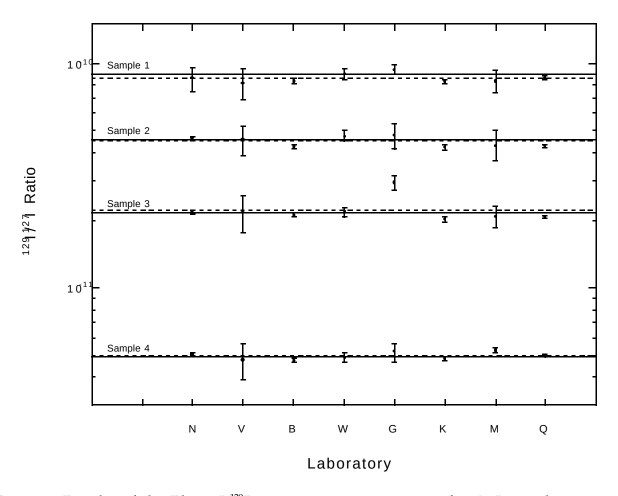


Figure 1. Results of the Phase I 129 I intercomparison exercise for AgI samples 1, 2, 3, and 4. The dashed line is an un-weighted mean of the results. The solid line is the expected value based upon successive dilution's of a NIST standard material. As one can see, the agreement between different laboratories for AgI is quite good.

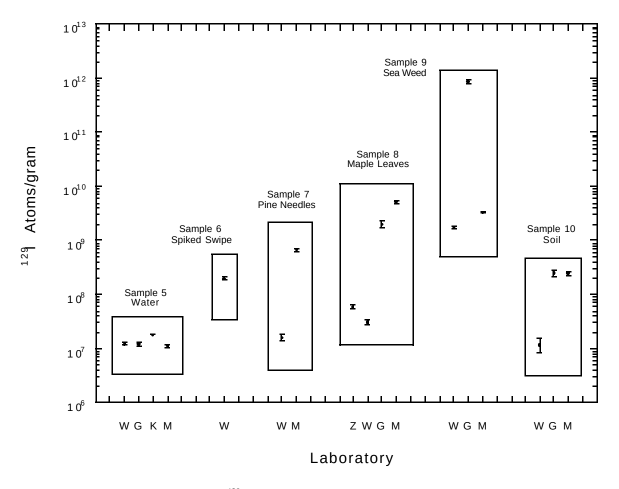


Figure 2. Results of the Phase I 129 I intercomparison exercise for samples 5, 6, 7, 8, 9, and 10. Except for sample 5, the agreements between measured 129 I concentrations are poor.

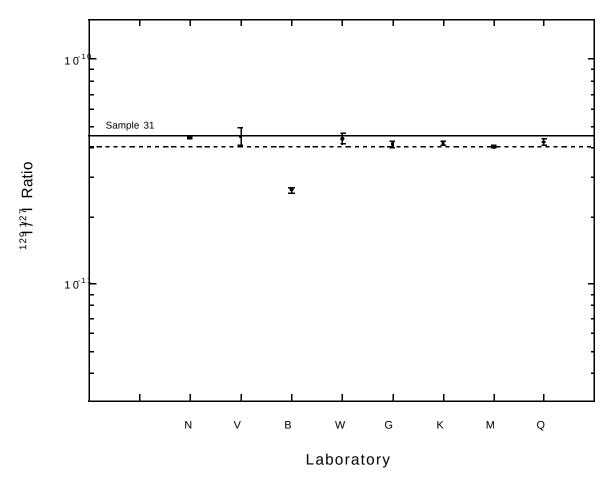


Figure 3. Results of the Phase II ¹²⁹I intercomparison exercise for AgI sample 31. Sample 31 is a repeat of sample 2 used in Phase I of the exercise. The dashed line is the un-weighted mean of the results. The solid line is the expected value based upon successive dilution's of a NIST standard material. With the exception of one point, the agreement between different laboratories for AgI is quite good.

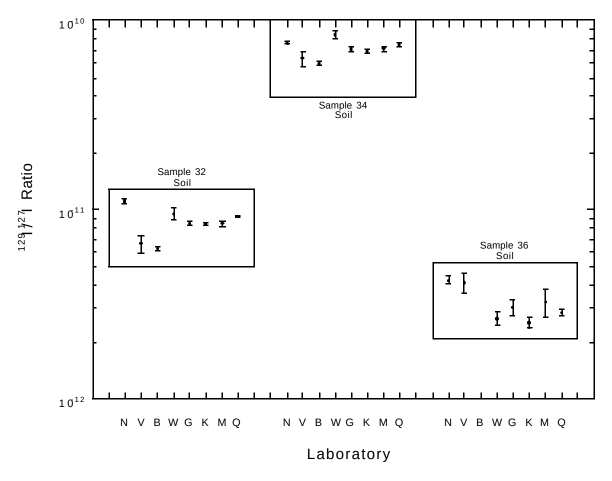


Figure 4. Measured ¹²⁹I/¹²⁷I ratios for samples 32, 34, and 36 of Phase II of the ¹²⁹I intercomparison exercise. Samples 32, 34, and 36 are AgI prepared from IAEA 375 soil using three differing chemical preparation methods. ¹²⁹I/¹²⁷I ratios for samples 32, 34, and 36 are not expected to agree because of the differing amount of carrier material used in the differing sample preparation methods.

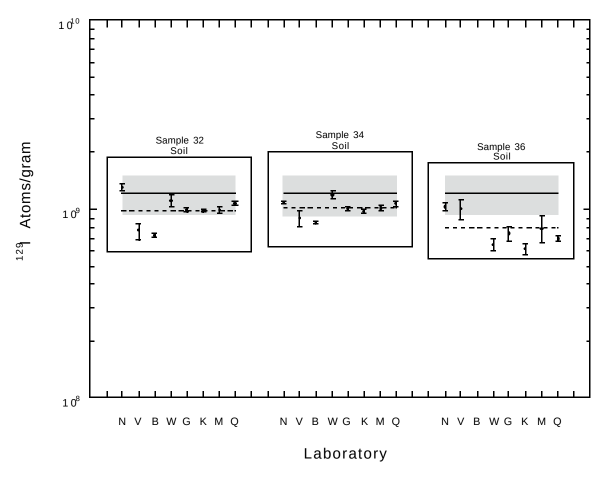


Figure 5. Measured 129 I concentrations for samples 32, 34, and 36 of Phase II of the 129 I intercomparison exercise. Samples 32, 34, and 36 were prepared from IAEA 375 soil using three differing chemical preparation methods. The dashed line is an un-weighted mean of the various results. The solid line is the reported IAEA 129 I concentration value of $1.7(\pm0.4) \times 10^{-3}$ Bq/kg or $1.2(\pm0.3) \times 10^{9}$ I atoms/gram material. The grayed area is the bounds the upper and lower limits of the IAEA reported value. As one can see, the agreement between differing measurements and differing chemical preparation methods is quite good.

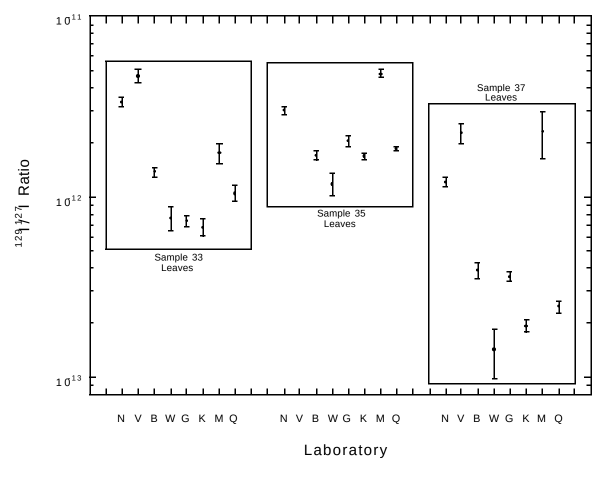


Figure 6. Measured ¹²⁹I/¹²⁷I ratios for samples 33, 35, and 37 of Phase II of the ¹²⁹I intercomparison exercise. Samples 33, 35, and 37 are AgI prepared from maple leaves using three differing chemical preparation methods. ¹²⁹I/¹²⁷I ratios for samples 33, 35, and 37 are not expected to agree because of the differing amount of carrier material used in the differing sample preparation methods.

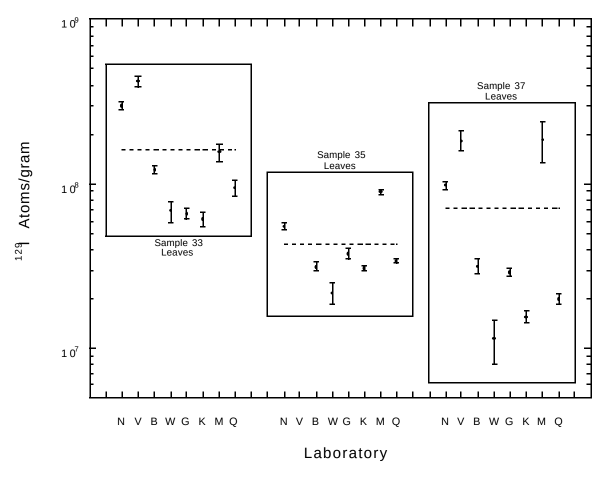


Figure 7. Measured ¹²⁹I concentrations for samples 33, 35, and 37 of Phase II of the ¹²⁹I intercomparison exercise. Samples 33, 35, and 37 were prepared from maple leaves using three differing chemical preparation methods. The dashed line is an un-weighted mean of the various results. As one can see, the agreement between differing measurements and differing chemical preparation methods is not very good.

(Results current as of 10/1/96)

Laboratory	ry Sample #1 Sample #		Sample #2	le #2 Sample #3			Sample #4	
Code	ž – – – – – – – – – – – – – – – – – – –		AgI		AgI		AgI	
	¹²⁹ I/ ¹²⁷ I Ratio		¹²⁹ I/ ¹²⁷ I Ratio		¹²⁹ I/ ¹²⁷ I Ratio		¹²⁹ I/ ¹²⁷ I Ratio	
N	8.61E-11 ± 1.05E-11		4.65E-11 ± 4.00E-13		2.17E-11 ± 4.00E-13		5.05E-12 ± 9.00E-14	
V	8.20E-11 ± 1.30E-11		4.60E-11 ± 7.00E-12		2.16E-11 ± 4.00E-12		4.80E-12 ± 9.00E-13	
В	8.41E-11 ± 2.30E-12		4.28E-11 ± 8.00E-13		2.13E-11 ± 4.00E-13		4.76E-12 ± 8.00E-14	
W	9.01E-11 ± 4.51E-12		4.78E-11 ± 2.39E-12		2.19E-11 ± 1.10E-12		4.90E-12 ± 2.45E-13	
G	9.40E-11 ± 5.00E-12		4.80E-11 ± 6.00E-12		2.95E-11 ± 2.40E-12		5.20E-12 ± 5.00E-13	
K	8.38E-11 ± 2.10E-12		4.22E-11 ± 1.03E-12		2.03E-11 ± 5.40E-13		4.85E-12 ± 1.20E-13	
M	8.39E-11 ± 9.70E-12		4.35E-11 ± 6.54E-12		2.09E-11 ± 2.28E-12		5.28E-12 ± 1.93E-13	
Q	8.66E-11 ± 1.50E-12		$4.32E-11 \pm 7.00E-13$		2.07E-11 ± 3.00E-13		5.00E-12 ± 7.00E-14	
Mean \pm Std. Dev.	8.63E-11 ± 3.95E-12	5%	4.50E-11 ± 2.34E-12	5%	2.22E-11 ± 2.98E-12	13%	4.98E-12 ± 1.88E-13	4%
Expected Ratio	9.03E-11		4.55E-11		2.17E-11		4.92E-12	
Laboratory	Sample #5		Sample #6		Sample #7		Sample #8	
Code Water Sample			Spiked Swipe 129I atoms/gram		Pine Needles	Pine Needles		
	¹²⁹ I atoms/gram				¹²⁹ I atoms/gram		¹²⁹ I atoms/gram	
Z							$5.89E+07 \pm 4.70E+06$	
В	$1.20E+07 \pm 3.00E+05$		Below Detection Limit		$1.10E+07 \pm 6.00E+05$		$2.50E+07 \pm 9.00E+06$	
W	$1.22E+07 \pm 6.10E+05$		$1.97E+08 \pm 1.38E+07$		$1.62E+07 \pm 2.43E+06$		$3.10E+07 \pm 4.65E+06$	
G	G $1.21E+07 \pm 8.00E+05$						$2.00E+09 \pm 3.00E+08$	
K	$1.83E+07 \pm 4.80E+05$							
M	$1.12E+07 \pm 4.61E+05$				$6.80E+08 \pm 5.18E+07$		5.21E+09 ± 1.92E+08	
Laboratory	Sample #9		Sample #10		Sample #11			
Code	Sea Weed		Soil		Woodward Iodine			
	¹²⁹ I atoms/gram		129I atoms/gram		¹²⁹ I/ ¹²⁷ I Ratio			
В			$2.40E+07 \pm 2.00E+06$					
W	$1.78E+09 \pm 1.25E+08$		$1.16E+07 \pm 3.48E+06$		$8.00E-14 \pm 2.00E-14$			
G	$8.70E+11 \pm 7.00E+10$		$2.50E+08 \pm 4.00E+07$					
K					9.00E-14 ± 3.00E-14			
M	$3.32E+09 \pm 1.29E+08$		$2.46E+08 \pm 1.38E+07$					
Q					5.30E-14 ± 1.60E-14			

Table 2
Results of Phase II of the ¹²⁹I Round Robin Exercise

(Results current as of 5/14/97)

Laboratory Code	Sample #31 AgI ¹²⁹ I/ ¹²⁷ I Ratio		Sample #32 AgI from Soil ¹²⁹ I/ ¹²⁷ I Ratio		Sample #33 AgI from Leaves 129I/127I Ratio		Sample #34 AgI from Soil ¹²⁹ I/ ¹²⁷ I Ratio	
N	4.47E-11 ± 6.50E-13		1.11E-11 ± 4.00E-13		3.36E-12 ± 2.00E-13		7.69E-11 ± 1.40E-12	
V	$4.54E-11 \pm 4.00E-12$		$6.61E-12 \pm 6.60E-13$		$4.71E-12 \pm 4.00E-13$		$6.31E-11 \pm 6.00E-12$	
В	2.62E-11 ± 6.59E-13		6.21E-12 ± 1.60E-13		$1.38E-12 \pm 8.00E-14$		5.95E-11 ± 1.39E-12	
W	$4.42E-11 \pm 2.21E-12$		9.54E-12 ± 6.68E-13		$7.66E-13 \pm 1.15E-13$		$8.44E-11 \pm 4.22E-12$	
G	$4.19E-11 \pm 1.50E-12$		8.50E-12 ± 1.90E-13		$7.40\text{E-}13 \pm 5.00\text{E-}14$		$7.03E-11 \pm 1.60E-12$	
K	$4.22E-11 \pm 8.40E-13$		8.44E-12 ± 1.90E-13		$6.83E-13 \pm 6.90E-14$		$6.87\text{E-}11 \pm 1.60\text{E-}12$	
M	$4.09E-11 \pm 6.80E-13$		$8.49E-12 \pm 2.80E-13$		$1.75E-12 \pm 2.10E-13$		$7.10\text{E-}11 \pm 2.50\text{E-}12$	
Q	$4.30E-11 \pm 1.40E-12$		9.18E-12 ± 1.50E-13		$1.05E-12 \pm 1.10E-13$		$7.45E-11 \pm 2.60E-12$	
Mean ± Std. Dev.	4.11E-11 ± 6.19E-12	15%	8.51E-12 ± 1.57E-12	18%	1.80E-12 ± 1.47E-12	81%	7.10E-11 ± 7.82E-12	11%
Expected Ratio	4.55E-11							

Laboratory Code	Sample #35 Sample #36 AgI from Leaves AgI from Soil $^{129}I/^{127}I$ Ratio $^{129}I/^{127}I$ Ratio						
N	3.03E-12 ± 1.40E-13		4.25E-12 ± 1.80E-13		1.21E-12 ±	7.00E-14	
V			4.12E-12 ± 5.00E-13		2.27E-12 ±	3.00E-13	
В	1.71E-12 ± 1.13E-13				3.91E-13 ±	3.90E-14	
W	1.18E-12 ± 1.77E-13		2.67E-12 ± 2.14E-13		$1.42\text{E-}13~\pm$	4.26E-14	
G	2.05E-12 ± 1.50E-13		$3.05E-12 \pm 3.00E-13$		$3.60\text{E-}13~\pm$	2.00E-14	
K	$1.68E-12 \pm 7.40E-14$		2.54E-12 ± 1.60E-13		$1.93\text{E-}13 \pm$	1.50E-14	
M	$4.83E-12 \pm 2.10E-13$		$3.25E-12 \pm 5.40E-13$		$2.29E-12 \pm$	6.50E-13	
Q	$1.85E-12 \pm 5.60E-14$		2.86E-12 ± 1.10E-13		$2.47E-13 \pm$	1.90E-14	
Mean ± Std. Dev.	2.33E-12 ± 1.24E-12	53%	3.25E-12 ± 6.82E-13	21%	8.88E-13 ±	9.22E-13	104%

 $Table\ 3$ Results of Phase II of the ^{129}I Round Robin Exercise

(Results current as of 5/14/97)

Laboratory	Sample #32	Sample #34	Sample #36		
Code	Soil	Soil	Soil		
	(atoms 129I/gram of material)	(atoms 129I/gram of material)	(atoms 129I/gram of material)		
N	$1.30E+09 \pm 4.66E+07$	$1.09E+09 \pm 1.99E+07$	$1.03E+09 \pm 4.37E+07$		
V	$7.69E+08 \pm 7.68E+07$	$8.96E+08 \pm 8.52E+07$	$1.00E+09 \pm 1.22E+08$		
В	$7.23E+08 \pm 1.86E+07$	$8.45E+08 \pm 1.97E+07$			
W	$1.11E+09 \pm 7.77E+07$	$1.20E+09 \pm 5.99E+07$	$6.49E+08 \pm 5.19E+07$		
G	$9.89E+08 \pm 2.21E+07$	$9.99E+08 \pm 2.27E+07$	$7.41E+08 \pm 7.29E+07$		
K	$9.82E+08 \pm 2.21E+07$	$9.76E+08 \pm 2.27E+07$	$6.17E+08 \pm 3.89E+07$		
M	$9.88E+08 \pm 3.26E+07$	$1.01E+09 \pm 3.55E+07$	$7.90E+08 \pm 1.31E+08$		
Q	$1.07E+09 \pm 1.75E+07$	$1.06E+09 \pm 3.69E+07$	$6.95E+08 \pm 2.67E+07$		
Mean ± Std. D	ev. 9.91E+08 ± 1.83E+08 18	1.01E+09 ± 1.11E+08	11% 7.90E+08 ± 1.66E+08 21%		

Laboratory	Laboratory Sample #33		Sample #35		Sample #37		
Code	Code Leaves		Leaves		Leaves		
	(atoms 129I/gram of material)) (a	toms 129I/gram of mate	erial)	(atoms 129I/gram of materia	al)	
N	$3.01E+08 \pm 1.79E+07$		5.57E+07 ± 2.57E+0	06	$9.80E+07 \pm 5.67E+06$		
V	$4.21E+08 \pm 3.58E+07$				$1.84E+08 \pm 2.43E+07$		
В	$1.23E+08 \pm 7.16E+06$		$3.14E+07 \pm 2.08E+0$	06	$3.17E+07 \pm 3.16E+06$		
W	$6.85E+07 \pm 1.03E+07$		$2.17E+07 \pm 3.25E+0$	06	$1.15E+07 \pm 3.45E+06$		
G	$6.62E+07 \pm 4.47E+06$		$3.77E+07 \pm 2.76E+0$	06	$2.92E+07 \pm 1.62E+06$		
K	$6.11E+07 \pm 6.17E+06$		$3.09E+07 \pm 1.36E+0$	06	$1.56E+07 \pm 1.22E+06$		
M	$1.57E+08 \pm 1.88E+07$		$8.88E+07 \pm 3.86E+0$	06	$1.86E+08 \pm 5.27E+07$		
Q	$9.40E+07 \pm 9.84E+06$		$3.40E+07 \pm 1.03E+0$	06	$2.00E+07 \pm 1.54E+06$		
Mean ± Std. De	ev. $1.62E+08 \pm 1.31E+08$	81%	4.29E+07 ± 2.27E+0	07 53%	$7.19E+07 \pm 7.47E+07$	104%	